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Non-methane hydrocarbons source apportionment at different sites in Mexico City during 2002–2003

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Abstract

The atmospheric concentrations of a variety of non-methane hydrocarbons (NMHC) collected at different sites, representing urban and rural environments within Mexico City Metropolitan Area (MCMA) during 1997, 2002 and 2003 field campaigns, were compared and used as an input for the Chemical Mass Balance (CMB) receptor model to determine the source contribution of NMHC to the atmosphere. A common feature at all the locations was the dominance of alkenes (59%), aromatics (16%) and olefins (9%) in the average NMHC burden. At the urban sites the interquartile range of NMHC concentrations showed stabilization over this period with a slight increase in the concentrations of propane and butanes in the southwest site of the MCMA in 2003 due to the increased use of liquefied petroleum gas (LPG). The receptor model CMB version 8.0 was used to apportion the NMHC sources at six locations within the MCMA, representing the heavily industrialized, commercial, residential and rural areas. For the 2003 field campaign, the contribution of vehicular emissions dominated the NMHC concentrations (19.7%±7.1% for gasoline vehicles and 35.4%±17.5% for diesel vehicles) followed by the emissions of marketing and handling of LPG (29.9%±8.0%).

The NMHC concentrations showed a weekly cycle with the highest levels towards the end of the week and lowest at weekend and beginning of the week, suggesting that both emissions and accumulations process play a key role in building up NMHC levels. The toluene to benzene ratio was used to determine photochemical ageing of the air samples during the 2003 field campaign. The database was divided into periods with similar wind circulation pattern; the results suggest that ageing process within the MCMA is generally suppressed by the amount of fresh emissions.

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1 Introduction

1.1 Background

Source based models have been used to identify and apportion non-methane hydrocarbons (NMHC) sources, which rely on the knowledge of emissions to predict their concentrations at different receptor sites (Scheff and Klevs, 1987; Li et al., 2003; Fujita et al., 1994; Vega et al., 1997 and 2000; Viana, et al., 2007). The Chemical Mass Balance (CMB) receptor model infers source contributions by determining the best fit combination of chemical profiles of emission source needed to reconstruct the chemical composition of ambient samples. This model is deterministic and does not rely on statistical interferences since it uses the chemical and physical characteristics of the gases (Watson et al., 1990, 1994; Core, 1987). It has been used in the analysis of suspended particles in the atmosphere (Watson et al., 1994; Vega et al., 1997) and was extended for the analysis of NMHC. The CMB model has also been used to evaluate the efficiency of specific control strategies associated with local air quality management programs and to estimate the emission inventory uncertainty (Scheff et al., 1984), since it correlates the pollutants with their sources of emission.

Since 1990 several air quality management programs and control strategies to reduce air pollution involving government, industry and inhabitants were implemented in Mexico City Metropolitan Area (MCMA) (Programa para Mejorar la Calidad del Aire en el Valle de Mexico 1995–2000) (Departamento del Distrito Federal, 1996). As a consequence, concentrations of sulfur dioxide, carbon monoxide and lead have been successfully controlled, but high ozone and particulate matter are still the major atmospheric pollution problem (Molina and Molina, 2002). Attention to the presence of atmospheric NMHC has been intensified because they are precursors to ozone and secondary organic aerosols and also because of their toxic and carcinogenic properties. To formulate and evaluate NMHC control strategies, the composition, location and concentration of these compounds must be determined.

The first part of this article presents the spatial and temporal analysis of the ambient

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concentrations of the most abundant NMHC which were measured during three field campaigns (1997, 2002 and 2003) at several monitoring sites within the MCMA (Molina et al., 2007). The photochemical age of the air masses was deduced by obtaining the toluene/benzene ratio, and the results were also analyzed in terms of the wind flow pattern. The second part of this article discusses the results of source identification and apportionment of NMHC to the atmosphere at four urban sites in the MCMA and two boundary sites within rural areas, applying the CMB receptor model.

1.2 CMB Receptor Model

The CMB modeling procedure requires: 1) identification of the contributing sources types; 2) selection of chemical species or other properties to be included in the calculation; 3) estimation of the fraction of each of the chemical species which is contained in each source type (source profiles); 4) estimation of the uncertainty in both ambient concentrations and source profiles; and 5) solution of the chemical mass balance equations. The CMB is implicit in all factor analysis and multiple linear regression models that intend to quantitatively estimate source contributions (Watson, 1984). The basic assumptions are: 1) composition of source emission are constant over the period of ambient and source sampling; 2) chemical species do not react with one another; 3) all sources which may significantly contribute to the receptor have been identified and their emissions characterized; 4) source profiles are linearly independent. The general equation is

$$C_i = \sum_{j=1}^P F_{ij} \cdot S_j$$

where C_i is the ambient concentration of the species “i” measured in the receptor site, P is the number of sources that contribute ($j = 1, 2, \dots, J$), F_{ij} is the fraction of the emissions of the species “i” starting from the source “j”, S_j is the impact to the receptor (calculated contribution) of the source “j”.

The parameters used to determine the performance of the model are: 1) R^2 (measures the variance in the receptor concentrations, which is explained by the calculated species concentrations, value 1); 2) χ^2 (is the weighted sum of the squared of differences between calculated and measured species divided by the effective variance and degrees of freedom; 3) percent of mass accounted for (is the ratio of the sum of the source contributions to the measured mass (Gertler et al., 1996). Calculations of the source contributions for Mexico City were carried out using CMB version 8.0 available from the U.S. EPA/Desert Research Institute.

2 Methods

2.1 Sampling sites

The Mexican Petroleum Institute (IMP) conducted two sampling campaigns to obtain the ambient data during March 2002 and April-May 2003 at different sites with sampling periods of 3 hr from 6:00 to 9:00 a.m (Velasco et al., 2007). The sampling sites encompass residential, industrial and mixed settings, and all of them classified as urban sites. The Northeast site, Xalostoc (XAL), is under the influence of heavily travelled paved and unpaved roads with old and new gasoline and diesel vehicles and historically has shown the highest concentrations of NMHC in Mexico City with major industrial activities. La Merced (MER) site is located in the administrative district in Central city, under the influence of commercial activities, residential and heavily traveled paved roads with light-duty vehicles and modern heavy-duty diesel buses. The southwest site is located in Pedregal (PED), in a residential area, with paved roads lightly travelled. The southeast site CENICA (CEN) is located near stone-cutting operations and heavily travelled paved and unpaved roads with old and new gasoline and diesel vehicles.

The Washington State University (WSU) conducted a sampling campaign to collect ambient air data during April 2003 at two boundary sites within rural areas (La Reforma and Santa Ana) (Velasco et al., 2007), and at three urban sites (La Merced, CENICA

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and Pedregal). The southwest site Santa Ana (SAN) is a rural site near Amecameca's Mountains surrounded by cactus fields and with paved and unpaved roads. The north-east site of La Reforma is a rural site located in the southwest of Pachuca (PAC), and close to urban areas with paved and unpaved roads.

5 2.2 Sampling

Sampling and analysis of ambient air and source emission profiles were carried out by IMP according to the Environmental Protection Agency (EPA) TO-14 method (Compendium method TO-14, 1988). The collection of ambient air samples was conducted using programmable ANDERSEN equipment (Arriaga et al., 1996), where sampling
10 period and starting time of sampling were programmable. The portable equipment has an electronic flow controller that allows the uptake of ambient air into the container to less than 25.0 psi. The feeding flow was adjusted prior to sampling to 66.6 ml/min. Sampling volumes were 6.0 litres taken throughout 3.0 hr. Samples for emission profiles were collected directly in the sources using canisters of 0.8 litres capacity. A flow
15 regulator of critical orifice was used to adjust the feeding flow between 80 to 13 ml/min and therefore, to control the sampling time (Vega et al., 2000). Samplings carried out by WSU of ambient air were collected by using a XonTech, Inc. Air Sampler model 910PC (Velasco et al., 2007).

2.3 Analysis of samples

20 Samples collected at XAL, MER, PED and CEN by IMP were analyzed at the IMP Atmospheric Chemistry Laboratory for NMHC (C2-C12 hydrocarbons). More than 200 compounds were determined although only a total of 49 species are reported, as they constitute more than 90% of the total NMHC mass. After cryogenic sample concentration in a freeze-out loop made from chromatographic grade stainless
25 steel tubing packed with 60/80 mesh deactivated glass beads, high-resolution capillary gas chromatography (GC) with flame ionization detector (Hewlett-Packard 5890

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Series II Plus) equipped with a 60 m Quadrex fused silica glass capillary column (0.32 i.d. and coated with a 1 μ m film thickness) at a flowrate of 2 ml/min. was used. The trap was then flushed-heated with boiling water and via a rotary valve (Six port Valco valve) switched to transfer the condensed NMHC into the GC equipment for analysis. The oven temperature started at -50°C and was increased to 200°C at a rate of $8^{\circ}\text{C}/\text{min}$. The chromatograph was calibrated by injecting known amounts of a mixture of 55 hydrocarbons (Scott Specialty Gases NIST Traceable), and a certified mixture of 33 halogen-containing compounds (Spectra Gases, with 10% analytical accuracy). The detection limit was determined to be 1 ppbC (Velasco et al., 2007).

Samples collected at XAL, MER, PED, CEN, PAC and SAN by researchers from WSU were analyzed by using a Hewlett-Packard 6890 Series chromatograph equipped with 30 m fused silica DB-1 column (0.32 i.d. and 1 μ m film thickness) with 2 ml/min carrier flow. Prior to sampling injection, the oven was cooled to -50°C . During analysis, the oven temperature was raised at $4^{\circ}\text{C}/\text{min}$ to a final temperature of 150°C . Detector response was calibrated with NIST traceable 2,2-dimethylbutane standard; the detection limit was determined to be 20 pptC (Velasco et al., 2007).

2.4 Source fingerprints

It should be noted that the major limitation to source identification is to distinguish between sources with similar emission source profile composition and also that a definitive emission profiles may not exist due to the complexity of the process itself, in particular combustion profiles. Further, vehicular emission profiles depend on numerous factors which may vary significantly from vehicle to vehicle using the same fossil fuel. Recently, Zavala et al. (2006) and Rogers et al. (2006) characterized real on-road emissions from specific types of vehicles using mobile laboratory operating in the “chase” mode during the MCMA-2002/2003 campaign and obtained mobile emissions of some of the hydrocarbons, while Schifter et al. (2003, 2004) estimated on road emissions of total hydrocarbons from remote sensing measurements.

Different NMHC source profiles were determined in Mexico City and further details

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about the source sampling have been previously described (Vega et al., 2000) and similar sampling procedures of sources were mentioned elsewhere (Vega et al., 2000b; Sánchez et al., 2001; Sánchez et al., 2004). The source profiles determined were: emissions of motor vehicle exhaust inside a tunnel and in dynamometer, hot soak measured in parking lots, emissions of motor vehicle exhaust using diesel at cross roads and inside parking lots, liquefied petroleum gas, asphalt plant, landfill, dry-cleaning using perchloroethylene, degreasing, paint used for vehicles, vinyl paints, varnishes used in furniture, graphic arts and food cooking processes (Vega et al., 2000). It should be mentioned that source profiles can be used as long as the process or the quality of each fuel type has not changed; the profiles are still representative of the emissions. Sources were selected according to their contribution of NMHC to the environment (Departamento del Distrito Federal, 1996). Figure 1 shows examples of source profiles used in this work.

2.5 Uncertainties of source profiles

Source categories for NMHC were identified by means of cluster analysis using STATISTICA v 5.1. The software was also used to calculate the variations between similar source profiles. Samples of each group were averaged to obtain the fraction of each NMHC in the profile. Uncertainties were calculated, as they are explicit inputs to the CMB model to calculate the standard error of the source contribution estimations and also to give less weight to those values that are near the detection limits. The uncertainties reported in this paper were calculated as the standard deviations from averaging the results obtained for each source profile and their replicates (Watson, 1984).

2.6 Fitting species and source categories

Main fitting species used for modelling were: propane, butane, i-butane, acetylene, pentane, hexane, toluene, xylenes, MTBE, 2,2,4,trimethylpentane, 2methylpentane and 3methylpentane since they are usually above minimum detectable limits (MDLs)

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and also because they are tracer of specific emission sources. The abundance of these species should be different at each source. This difference is used by the model to apportion the contribution of emission sources at a specific receptor site. A total of 33 source profiles were grouped into six source categories to aid in the interpretation of results: 1) Liquefied petroleum gas (LPG), 2) degreasing processes using toluene, m,p-xylenes, pentane and hexane (Degrease), 3) solvents used in paint, offset and graphic arts (Paint), 4) light-duty vehicle exhaust emissions, which were determined in tunnels, crossroads and dynamometer (Gasoline), 5) heavy-duty vehicle exhaust emissions, which were determined in tunnels, crossroad and parking lots (Diesel) and 6) non-identified sources (Other).

3 Results

3.1 Ambient concentrations of NMHC during 1997, 2002 and 2003 field campaigns

The atmospheric levels of NMHC (ppbC) measured in the three field campaigns are described in Fig. 2 which contains the descriptive statistics for the sites and year. The wind roses are included to aid in the discussion of results in the following paragraphs. Box-whisker plots on left side of Fig. 2 shows the results at the industrial site XAL, the NMHC median concentrations have shown small changes throughout time, fluctuating at around 2600 ppbC. The major changes were observed in the maximum levels which have decreased by about 1000 ppbC from 1997 to 2003. The site representing the commercial-residential zone MER, showed a clear reduction in the maximum levels (by more than 3000 ppbC) from 1997 to 2002. As Fig. 2 shows, the highest concentrations (max=6677 ppbC, P_{75} =5600 ppbC and P_{50} =3900 ppbC) were measured at this site in 1997. The subsequent campaigns showed no major changes on the median concentrations, stabilizing at around 2500 ppbC. The PED site has been characterized by having the lowest concentrations of NMHC, as it has no major sources of pollution other than vehicular emissions (Vega, et al., 2000). The descriptive statistics show

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that the levels of NMHC at this site had minor changes from 1997 to 2002 and a slight increase in 2003, which was driven by the increment of propane and i-butane (Fig. 3). Regardless of the differences that each site presented, the 25–75 percentile range has narrowed indicating that atmospheric levels of NMHC are stabilizing throughout time, with median concentrations around 2500 ppbC at XAL and MER, and 1300 ppbC at PED. These results agree with Velasco et al. (2007), who reported that ambient levels of NMHC are stabilizing in the MCMA, despite the growth in vehicular fleet.

3.1.1 Relationship between NMHC maximum concentration and wind circulation

A series of wind roses were obtained for the days when the maximum or minimum levels of NMHC were measured. In Fig. 2, upper/bottom wind roses correspond to the day of maximum/minimum NMHC levels, respectively. The prevalent wind direction at XAL indicated that NE winds were associated with maximum NMHC concentration, showing a slight variation during the 2003 field campaign. Northern winds were associated to minimum NMHC concentrations except for the 2003 campaign which showed winds blowing from the SW. The wind roses for the MER site showed high variation, particularly for the maximum NMHC concentration (6677 ppbC) in 1997. The subsequent campaigns showed winds blowing from the N or the E associated to the maximum NMHC. The SW sector was related to maximum NMHC at the residential site PED. The effect of wind speed on the NMHC levels was not clear other than the association of highest concentrations and winds below or about 1.0 m/s. De Foy et al. (2005) mentioned that early morning winds in the MCMA were generally below 2.0 m/s, and no clear association was found between wind direction and speed.

The wind circulation patterns during MCMA-2003 campaign has been studied on detail by de Foy et al. (2005) using both synoptic and basin observations, and their effect on ozone peak concentration. Three circulation patterns were defined under which ozone levels will be highest in the north of the city (O₃-North category), in the south of the city (O₃-South category) and a third flow pattern (Cold Surge category) which will lead to lower temperatures and presence of rain with a more even geographical distribution.

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bution of pollutant concentrations. De Foy et al. (2005) also reported that the frequency of ozone concentrations above 200 ppb was higher during the O₃-north flow, a lower but broad distribution of concentration for a Cold surge category and a narrow concentration around 170 ppb for an O₃-South category. These flow patterns described by de Foy also fitted to NMHC levels, as maximum concentrations at XAL and MER corresponded to the O₃-North pattern, and maximum NMHC at PED was observed during an O₃-South flow pattern, reflecting the coupled chemistry of ozone and NMHC.

3.1.2 Concentrations of NMHC and the ratio toluene/benzene during the 2003 field campaign

A general feature for the three field campaigns was the observation of maximum concentrations of NMHC during weekdays especially from Wednesday to Friday, and minimum concentrations on weekends or at the beginning of the week (Monday or Tuesday). The variation of NMHC concentrations from Monday to Sunday is shown in Fig. 4, which was constructed using data from the six monitoring sites. These results suggest that NMHC concentration has a weekly cycle with the highest levels at the end of the week and lowest concentrations at weekend/beginning of week, resulting from both the emissions and accumulation process, the last one strongly dependent on meteorology.

The photochemical age of an air sample has been estimated by using ratios of measured NMHC (McKeen and Liu, 1993; Cubison et al., 2006). This method has a simple principle: at the emission source, NMHC have a characteristic ratio which depends on the type of emissions. Downwind emission source ratios will change as the more reactive species will be removed faster. In this work we use the toluene/benzene ratio as a method of estimating photochemical age (de Gouw et al., 2005). We used the data from source profiles studies in Mexico City (Vega et al., 2000) to derive emission ratios of vehicular emissions as they are the principal pollution source in urban environments, the results are presented in Table 1. MCMA-2003 campaign measurements were divided into periods depending on circulation flow pattern as described by de Foy et al. (2005), and only those measurements taken during the morning were used for

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this analysis.

Table 2 contains the date, type of day, concentration of NMHC, toluene/benzene ratio and wind circulation category. The first period from the 14th to 17th April showed an O₃-south pattern. The toluene/benzene ratio for sites located in the south of the MCMA (PED and SAN) showed an increase in the ratio after a 3-day period of O₃-South circulation pattern; besides, the maximum concentration of NMHC at PED (16th of April) was also observed. These ratios may reflect the contribution of NMHC sources as the air masses passed over the city, and the accumulation of pollutants in the south of the city. A Cold Surge O₃-South flow pattern started on the 19th of April showing ratios of 3.13 and 3.57 on the 21st and 22nd at MER site respectively, i.e., negligible change. An O₃-North flow pattern started on the 23rd through the 30th of April with a weekend in the middle of this period. The ratios at XAL increased from 4.35 on 23 April 2003 (Wednesday), 4.55 on 24 April 2003 and 5.88 on 25 April 2003. The ratios for 26 April 2003 (Saturday) and for 28 April 2003 showed a decrease. Actually, this was the only period when an ageing was observed. The results suggest that ambient air in MCMA is generally receiving fresh emissions high enough to avoid significant photochemical ageing, and depending on meteorology NMHC will accumulate in the north or in the south of the city. Velasco et al. (2007) found negligible photochemical ageing using ratios obtained from the diurnal profiles at CEN site. Figure 5 describes the relationship between NMHC concentration and toluene/benzene ratio divided by type of day. Although the coefficients were moderated, they show a negative relationship during weekends and positive but poor correlation during weekdays, supporting the assumption that it is possible to observe ageing process only during weekends (with lower emissions).

Figure 6 shows the variation on the toluene/benzene ratio for the three wind circulation categories. The highest ratios were observed during the O₃-North episodes, reflecting the higher amount of emissions and the accumulation process previously described.

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3.2 Source Apportionment results

Table 3 shows the average statistical parameters of the CMB model applied to NMHC at different sites within the MCMA during 2002–2003 field campaigns. In general, it can be observed that the parameters of R^2 , Chi^2 and percentage mass were within the acceptable range. Values of R^2 ranged from 0.80 at CEN to 0.99 at PED, MER and XAL, with an average of 0.95. In most cases, the values of Chi^2 were smaller than 4.00 with an average of 2.25. The percentages of mass calculated when applying the model varied from 80.3% at Pedregal to 114.4% at CEN, with an average of 91.3%. The results of measured and calculated ambient concentrations in ppbC when applying the model are also shown. It should be noted that in few cases, differences of calculated and measured mass were significant especially at the rural sites where mass was low, which indicates that there were species in the atmosphere that were not identified, this may be due to the location of diverse sources that contributed to ambient emissions but were not characterized.

3.2.1 Measurements from 06:00 to 09:00 and 12:00 to 15:00 during 2002 and 2003

In general, the results showed that the major sources of NMHC to the atmosphere were emissions from vehicle exhaust using gasoline, heavy duty vehicles using diesel and liquefied petroleum gas for both years as seen in Figs. 7 and 8. The main source at XAL was gasoline with an average of 40.6% in 2002, and diesel with an average of 47.4% in 2003, the average contribution of LPG was similar for both years. The main source at MER was gasoline with an average of 42.8% in 2002, and diesel with an average of 32.8% in 2003. The main source at PED was gasoline with an average of 41.0% in 2002, and LPG with an average of 32.8% in 2003, which is in agreement with the observed increment in propane and butane that will be discussed later.

A series of less than 60-min sampling measurements were conducted at MER, CEN and PED for some of the days during the 2003 field campaign. Some differences were observed when compared to the typical 3-hr sampling measurements on both

total NMHC concentration and source apportionment results. Such differences are described in the following paragraphs.

Figure 9A depicts the source apportionment results at MER site for the 23rd, 26th, 27th and 28th of April. Large differences were observed between the morning (06:00 to 09:00 hr) and the afternoon (16:00 to 19:00 hr) results, as the maximum contribution of LPG and gasoline vehicles occurred in the morning (07:00 to 08:00 hr), and minimum contribution in the afternoon, when the main source contribution shifted to diesel vehicles. As Figure A shows, the total NMHC concentration during the morning was 4 to 5-times higher than the afternoon values. A general feature was the higher concentration of total NMHC in the short-sampling measurements compared to the 3-hr samples, as more emissions from a specific point source can be captured when samples are taken on shorter periods of time. On the contrary, higher concentrations are missed in the 3-hr samples. This is in agreement with model results, in which smaller number of sources contributed to the 3-hr samples compared to the short-time samples. Figure 10A describes the source apportionment results for CEN site on the 8th, 9th, 13th, 14th of April, showing the contribution of sources which generally were not observed in the 3-hr sampling measurements (Fig. 8), although their contribution was not significant. These sources were degreasing and application of paint offset and graphic arts processes, all of them with emissions rich on toluene, xylenes and pentane. The LPG contribution, one of the major contributing sources, was constant (38%) for both short and 3-hr sampling measurements. The vehicular sources showed a variable contribution, i. e., the contribution of diesel vehicles was higher (58%) in the 3-hr sampling measurements compared to the short-sampling measurements (37.5% in average). The opposite was observed for the gasoline vehicles that showed a higher contribution from the short-sampling measurements (17.9%) compared to the 3-hr measurements (5.2%).

A comparison of short-sampling measurements vs. 3-hr sampling measurements at MER site showed that the main sources were LPG and diesel vehicles in both cases. The contribution of gasoline vehicles was higher in the short-sampling measurements

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(36% on average) than the 3-hr samples (24.8%). The contribution of printing processes was variable with an average of 9.8%.

Figure 9B shows the source apportionment for instantaneous sampling measurements at MER site, which were taken at 06:00, 07:00 and 08:00 hr. The total mass concentration was higher at 08:00 hr which is consistent with the short-sampling measurements characteristic. The main contribution sources were diesel vehicles with an average of 41.2%, LPG (29%) and gasoline vehicles (17.4%). It is important to mention that the non-identified sources showed their highest contribution 12.3% for this type of samples.

The short-sampling measurements at CEN on the 27th and 28th of April (Figs. 10B and 10C) indicated that higher concentrations were recorded at 07:30 and 08:00 hr, with a general decrease as the day goes on. It is worth mentioning that the model showed some difficulties apportioning the sources for the cases with high NMHC mass concentration, hence the contribution of non-identified sources increased. A slightly different pattern was observed on the 29th of April (Fig. 10D) that showed the maximum concentration at 08:00–08:30 hr. The contribution of diesel vehicles was homogeneous (36.5% on average). The LPG contribution was also important (37.5% on average) although showed some variation. The third most important source, gasoline vehicles, showed a range of contribution from 16.4% to 24.5% with an average of 19.5%.

Figure 9C describes the short-sampling measurements at PED site from 19th to 23rd of April. The main contribution was LPG with 43.8%, which is slightly higher than the 3-hr samples (37.8%); the second most important contribution shifted between gasoline vehicles (17.5% for the short-samples vs. 24.5% for the 3 hr samples) and diesel vehicles (29.8% for the short-samples vs. 21.7% for the 3-hr samples). The LPG contribution reached its highest contribution (up to 62% from 07:00 to 08:00 hr) at this site, which is consistent with the high observed concentrations of propane and butane.

The 3-hr measurements at the rural sites showed low concentrations, especially at Santa Ana site. The model results indicated that the main source was diesel vehicles

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(42.1% on average); gasoline vehicles reached 16.6% and LPG 15.9%. The contribution of non-identified sources was significant at both sites (25.5% on average) and was directly associated with the total NMHC concentration (Fig. 11).

Finally, a series of samples taken from 12:00 to 15:00 hr were analysed. The results indicated that source contribution was homogeneously distributed except for two days, when concentrations were higher than 1000 ppbC, although the total mass was not fully apportioned. In general, the contribution of LPG was found to be lower during the period from 12:00 to 15:00 hr compared to morning results. On the contrary, vehicular emissions increased during this period; the average contribution was 12.1% for LPG, 17.9% for gasoline and 47.7% for diesel (Fig. 12).

4 Conclusions

Non-methane hydrocarbon concentrations collected at different sites, representing urban and rural environments within the MCMA during 1997, 2002 and 2003 field campaigns, were compared and used as an input for the CMB model to determine the source contribution of NMHC to the atmosphere. For the 2003 field campaign, the results show that the contribution of vehicular emissions dominated the NMHC concentrations (19.7%±7.1% for gasoline vehicles and 35.4%±17.5% for diesel vehicles) followed by the emissions of marketing and handling of LPG (29.9%±8.0%). The latter source showed its higher contribution at the southwest site, especially during 2003 field campaign.

Despite the growth in the vehicular fleet, the observed concentrations of NMHC in several sites within the MCMA have shown stabilization over this period, as the 25–75 percentile range has narrowed, with median concentrations of 2500 ppbC at XAL and MER, and 1300 ppbC at PED in 2003. However, a slight increase in the concentrations of propane and i-butane was observed at the southwest site on 2003, which was attributed to a rise in the contribution of LPG emissions.

The effect of meteorological conditions over the NMCH spatial distribution was ana-

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lyzed using the 2003 field campaign data. It was observed that maximum concentrations of NMHC at XAL and MER took place during an O₃-north episode, and maximum NMHC at PED was observed during an O₃-south episode, reflecting the coupled chemistry of ozone and NMHC.

The NMHC concentrations showed a weekly cycle with the highest levels at the end of the week and lowest concentrations at weekend/beginning of week, suggesting that both emissions and accumulation process play a key role in building up NMHC levels. The toluene/benzene ratio of the air samples was calculated as a method for elucidating the photochemical age of the air samples. The ratios were analyzed as a function of the type of day, concentration of NMHC and wind circulation patterns. We found that the atmosphere of the MCMA generally receives fresh emissions high enough to prevent photochemical ageing. In addition, depending on meteorological conditions, NMHC will accumulate in the north or in the South of the city, increasing the toluene/benzene ratio after two or more days with similar flow pattern.

A series of instantaneous, 30-min and 60-min NMHC sampling were carried out for some days during the MCMA-2003 campaign. The results indicated that a more frequent sampling period (30 min) with respect to 3-h samplings showed higher concentrations since the longer sampling periods masked important NMHC variations, therefore, exposure to high levels could be missed. Some monitoring sites are influenced by nearby sources and represent neighborhood rather than specific local site exposure. For example, the CEN site was located near sources that contribute to the emission of NMHC to atmospheric loadings; XAL is influenced by emission from different industries and unpaved roads. These sources have a regional as well as local effect, but they may overestimate the total NMHC mass concentrations measured when the sampler is very close to the emission sources and may bias exposure estimates.

In summary, the spatial and temporal analysis of the ambient concentrations of the most abundant NMHC measured during three field campaigns from 1997 to 2003 was presented. The CMB receptor model was used to apportion the NMHC sources at six locations within the MCMA representing the heavily industrialized, commercial, resi-

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dential and rural areas. It should be noted that the sampling campaign was spatially and temporally limited and the data may reflect meteorological and other conditions specific to the sampling period. Therefore conclusions of this paper are specific to the sampling periods and locations in 2002 and 2003, and their applicability to other time periods and at other locations are not known. We are planning to apply similar technique to the datasets obtained from the MILAGRO (Megacity Initiative: Local and Global Research Observations) Campaign, a multi-component international scientific collaboration that took place in 2006, which will provide additional insights to the contribution of various sources to the NMHC burden.

Acknowledgements. We would like to thank the Washington State University and the staff of the Laboratorio de Química de la Atmósfera at the Instituto Mexicano del Petroleo for providing the data from analyses of the samples collected in the MCMA-2002/2003 campaigns. LTM acknowledges support from Mexican Comisión Ambiental Metropolitana and US National Science Foundation (ATM-0528227).

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Table 1. Toluene to benzene ratio for different emission sources in Mexico City (Vega, et al. 2000).

Source	Toluene/benzene
Whole gasoline	2.86
Hot soak emissions	1.68
Tunnel (gasoline vehicles)	2.45
Diesel vehicles	5.20

Table 2. Concentrations of NMHC, toluene to benzene ratio and flow category according to de Foy et al., (2005). Only morning samples are shown.

Date	Day	flow category	VOC conc (ppbC)	site/sample duration	Tol/ben
14 Apr 2003	Monday	O3-south	150	PAC 07-10	2.13
14 Apr 2003	Monday	O3-south	1176	PED 06-09	2.56
15 Apr 2003	Tuesday	O3-south	888	PAC 07-10	3.23
15 Apr 2003	Tuesday	O3-south	68	SAN 06-09	3.13
16 Apr 2003	Wednesday	O3-south	99	SAN 06-09	5.26
16 Apr 2003	Wednesday	O3-south	1951	PED 06-09	5.00
21 Apr 2003	Monday	cold surge	1252	MER 06-09	3.13
21 Apr 2003	Monday	cold surge	1356	CEN 06-09	3.13
21 Apr 2003	Monday	cold surge	1701	XAL 06-09	6.67
22 Apr 2003	Tuesday	cold surge	1189	MER 06-09	3.57
22 Apr 2003	Tuesday	cold surge	658	CEN 06-09	3.23
22 Apr 2003	Tuesday	cold surge	935	PED 06-09	3.45
23 Apr 2003	Wednesday	O3-north	3157	MER 06-09	9.09
23 Apr 2003	Wednesday	O3-north	2047	CEN 06-09	5.56
23 Apr 2003	Wednesday	O3-north	3367	XAL 06-09	4.35
23 Apr 2003	Wednesday	O3-north	1173	PED 06-09	3.23
24 Apr 2003	Thursday	O3-north	3036	MER 06-09	4.55
24 Apr 2003	Thursday	O3-north	2714	CEN 06-09	5.56
24 Apr 2003	Thursday	O3-north	4133	XAL 06-09	4.55
24 Apr 2003	Thursday	O3-north	953	PED 06-09	4.55
25 Apr 2003	Friday	O3-north	2490	MER 06-09	4.35
25 Apr 2003	Friday	O3-north	2245	CEN 06-09	6.25
25 Apr 2003	Friday	O3-north	2271	XAL 06-09	5.88
25 Apr 2003	Friday	O3-north	1425	PED 06-09	4.17
26 Apr 2003	Saturday	O3-north	2360	MER 06-09	4.76
26 Apr 2003	Saturday	O3-north	2057	CEN 06-09	11.11
26 Apr 2003	Saturday	O3-north	2580	XAL 06-09	4.35
26 Apr 2003	Saturday	O3-north	1560	PED 06-09	4.76
28 Apr 2003	Monday	O3-north	2300	MER 06-09	4.00
28 Apr 2003	Monday	O3-north	3605	CEN 06-09	5.00
28 Apr 2003	Monday	O3-north	3735	XAL 06-09	3.13
28 Apr 2003	Monday	O3-north	1549	PED 06-09	3.45
30 Apr 2003	Wednesday	O3-north	1751	MER 06-09	4.35
30 Apr 2003	Wednesday	O3-north	2700	CEN 06-09	4.76
30 Apr 2003	Wednesday	O3-north	2409	XAL 06-09	10.00
30 Apr 2003	Wednesday	O3-north	1400	PED 06-09	4.17
2 May 2003	Friday	O3-south	2204	MER 06-09	3.45
2 May 2003	Friday	O3-south	2032	CEN 06-09	3.85
2 May 2003	Friday	O3-south	2511	XAL 06-09	9.09
2 May 2003	Friday	O3-south	1056	PED 06-09	3.13

Table 3. Average statistical parameters of the CMB model applied to NMHC during 2002–2003 at different sites in Mexico.

Site		R ²	CHI ²	Meas. Conc. [ppbC]	Calc. Conc. [ppbC]	%Mass
Xalostoc (XAL)	Ave	0.97	2.04	2552	2280	89
	Min	0.87	0.91	719	675	80
	Max	0.99	3.63	4075	3882	100
Merced (MER)	Ave	0.96	2.14	2060	1937	94
	Min	0.89	1.01	366	328	81
	Max	0.99	3.87	3774	3626	113
Pedregal (PED)	Ave	0.95	2.53	839	748	89
	Min	0.86	0.98	320	293	80
	Max	0.99	4.07	1747	1592	104
Cenica (CEN)	Ave	0.94	2.16	1361	1281	93
	Min	0.80	0.88	253	204	80
	Max	0.98	3.75	4125	3877	114
Pachuca (PAC)	Ave	0.92	2.29	179	153	86
	Min	0.86	1.49	73	61	82
	Max	0.96	3.55	411	337	93
Santa Ana (SAN)	Ave	0.93	1.89	185	165	89
	Min	0.88	0.63	87	72	81
	Max	0.98	2.76	239	210	101
Spatial	Ave	0.95	2.24	1462	1348	91
	Min	0.8	0.63	73	61	80
	Max	0.99	4.07	4125	3882	114

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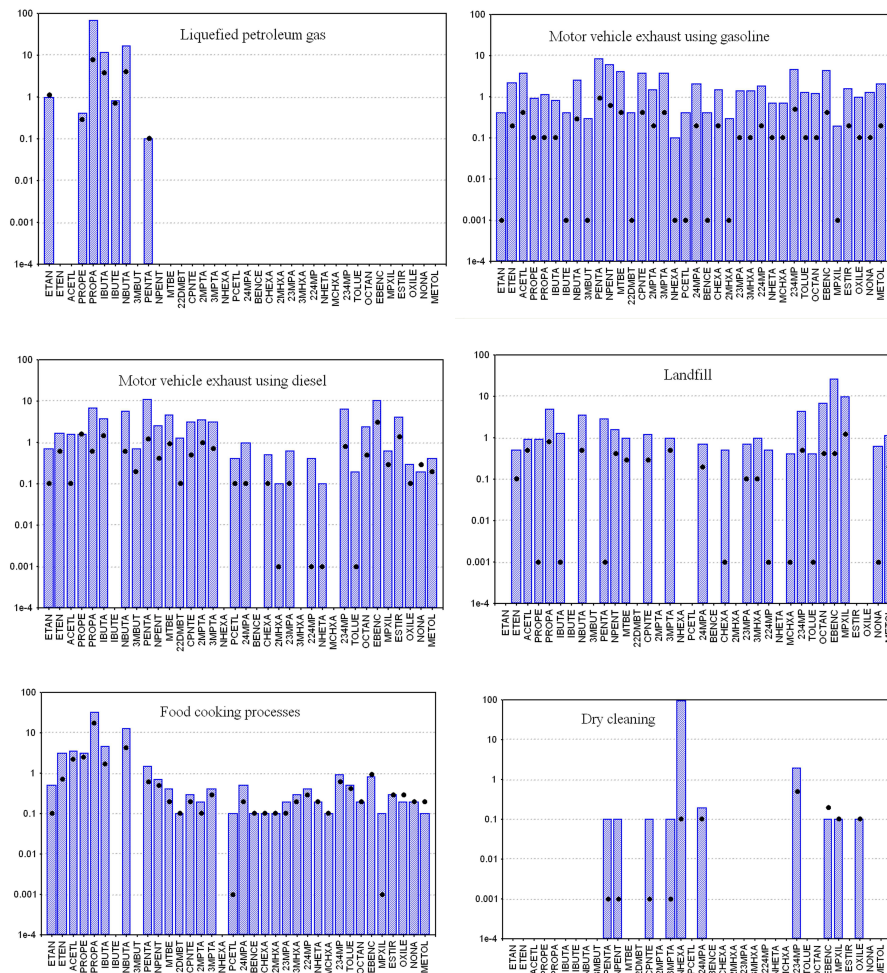


Fig. 1. Examples of NMHC source profiles used in the CMB model.

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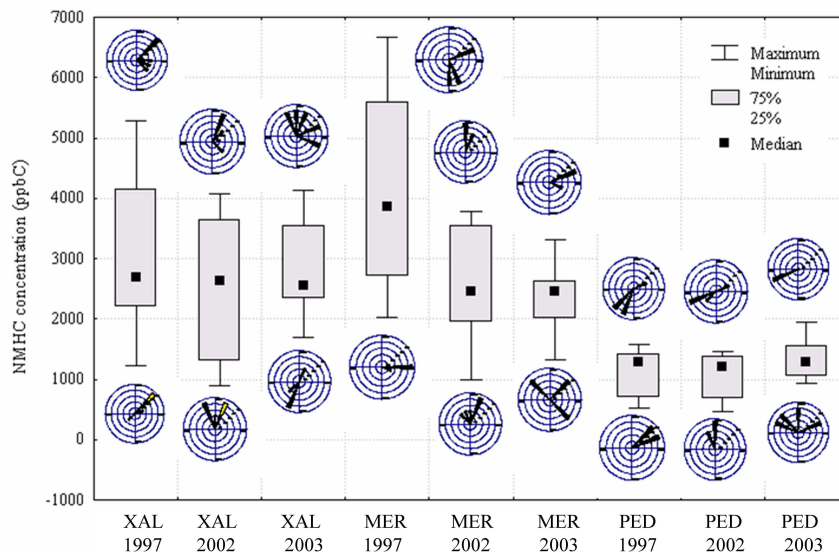


Fig. 2. Descriptive statistics for the NMHC concentrations (ppbC) measured at Xalostoc (XAL), La Merced (MER) and Pedregal (PED) during 1997, 2002 and 2003 field campaigns. The upper/bottom wind roses correspond to the day of maximum/minimum NMHC concentration, respectively.

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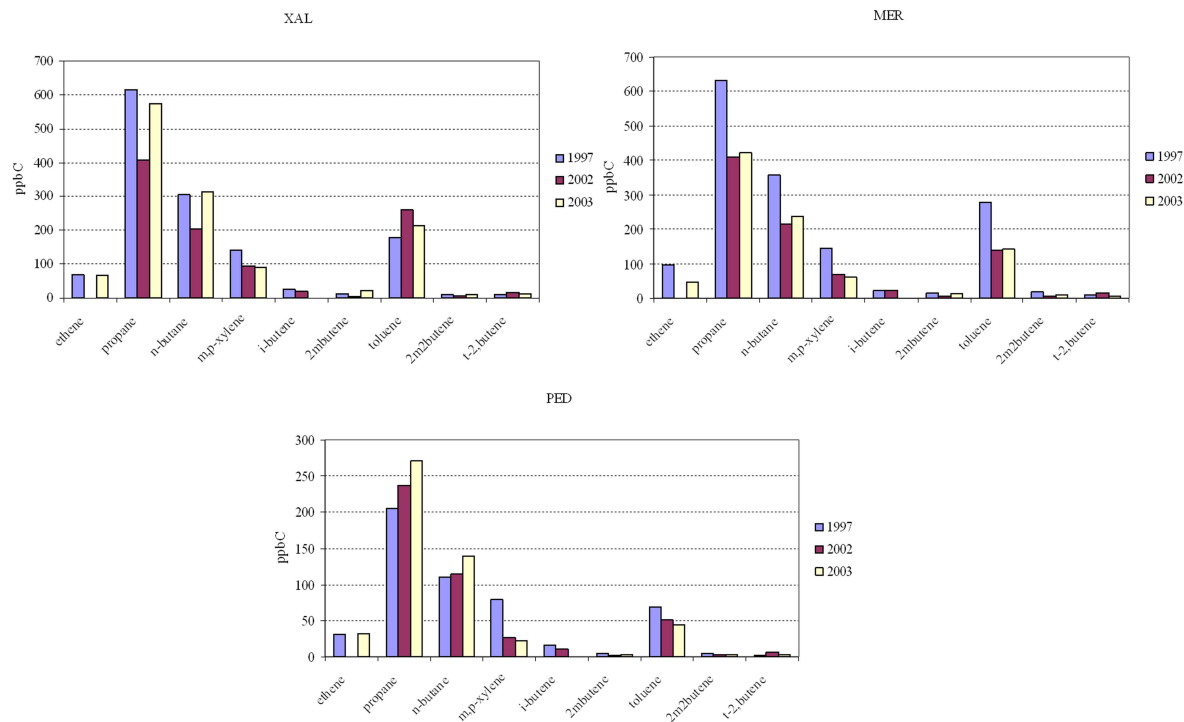


Fig. 3. Atmospheric concentration of the most abundant NMHC (ppbC) in the MCMA during the 1997, 2002 and 2003 field campaigns.

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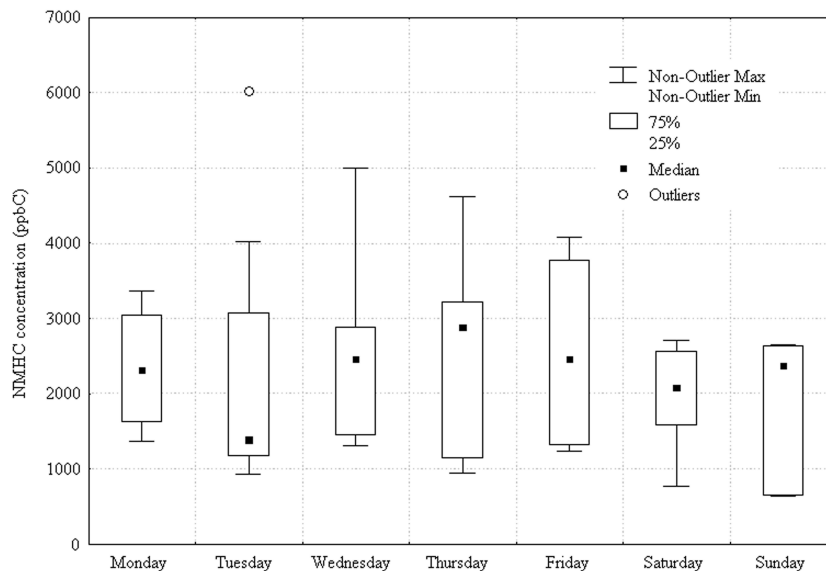


Fig. 4. Descriptive statistics for NMHC atmospheric concentrations (ppbC) in the MCMA grouped by type of day.

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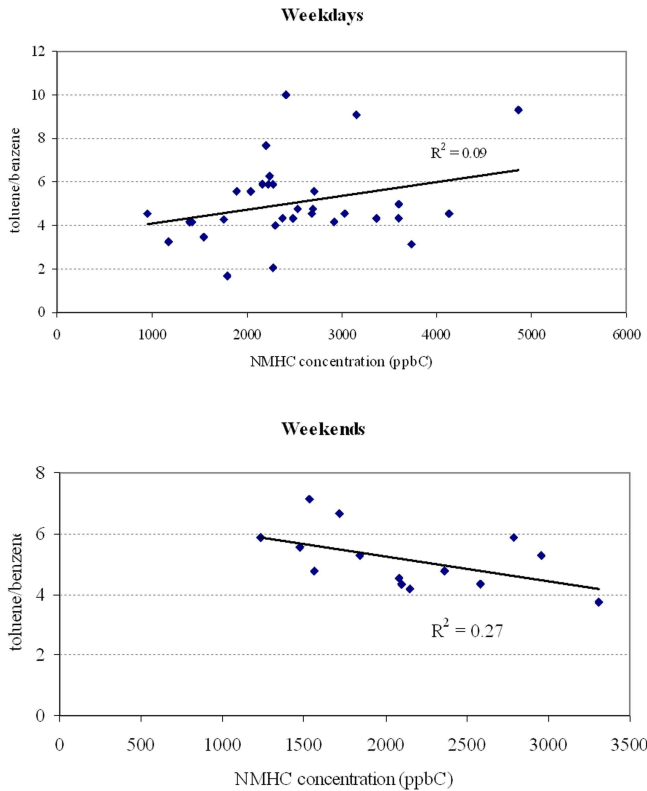


Fig. 5. Scatter plot between toluene/benzene ratio and NMHC concentration for weekends and weekdays.

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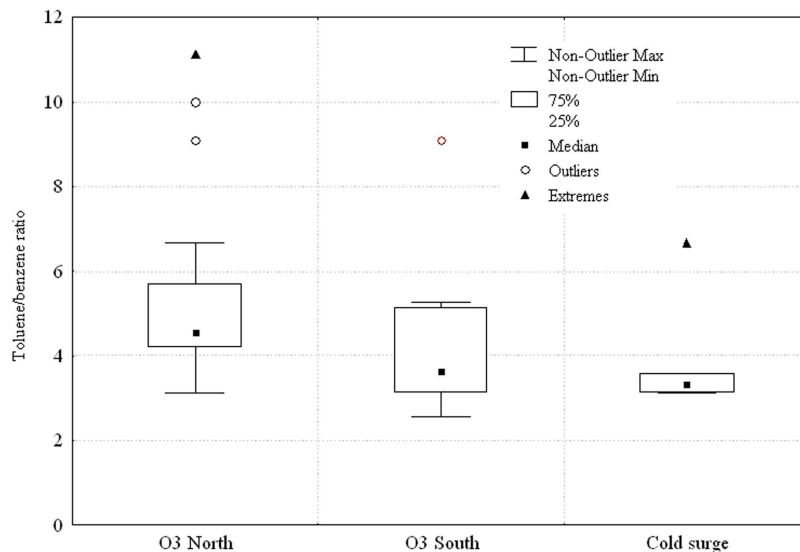


Fig. 6. Descriptive statistics for toluene/benzene ratio divided into flow categories during the 2003 field campaign.

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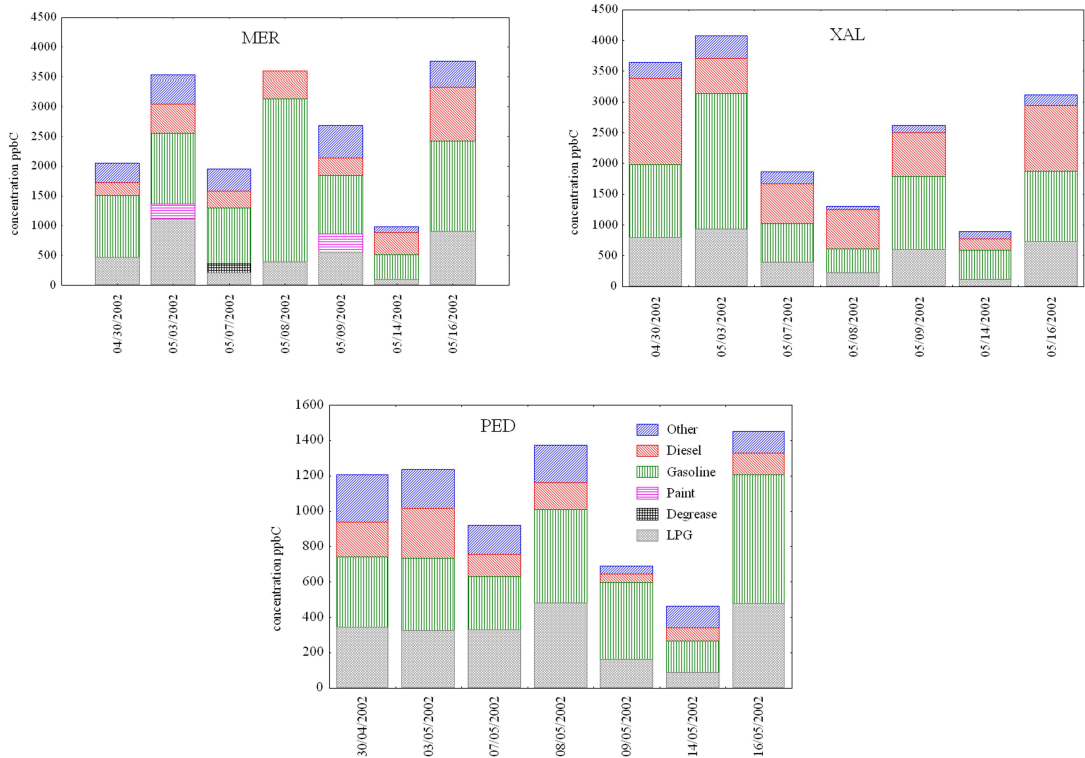


Fig. 7. Source apportionment distribution for NMHC during the 2002 campaign. 06:00–09:00 hr.

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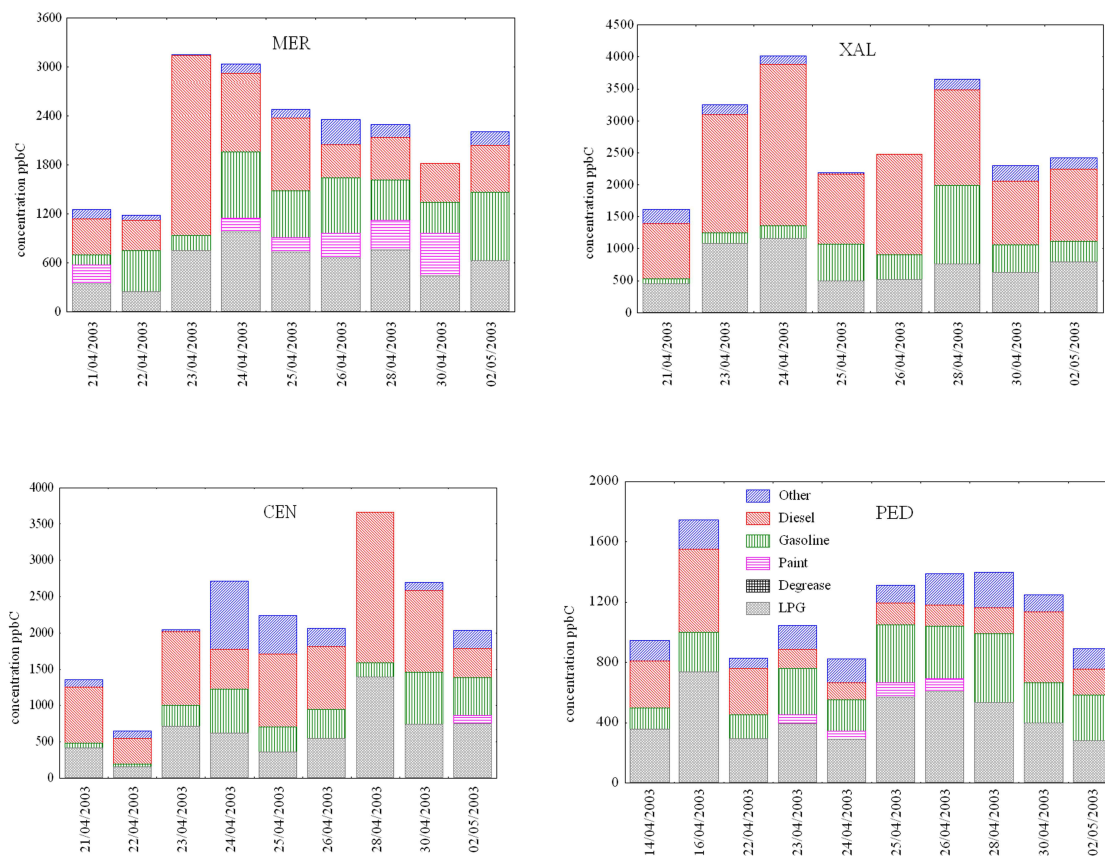


Fig. 8. Source apportionment distribution for NMHC during the 2003 campaign. 06:00–09:00 hr

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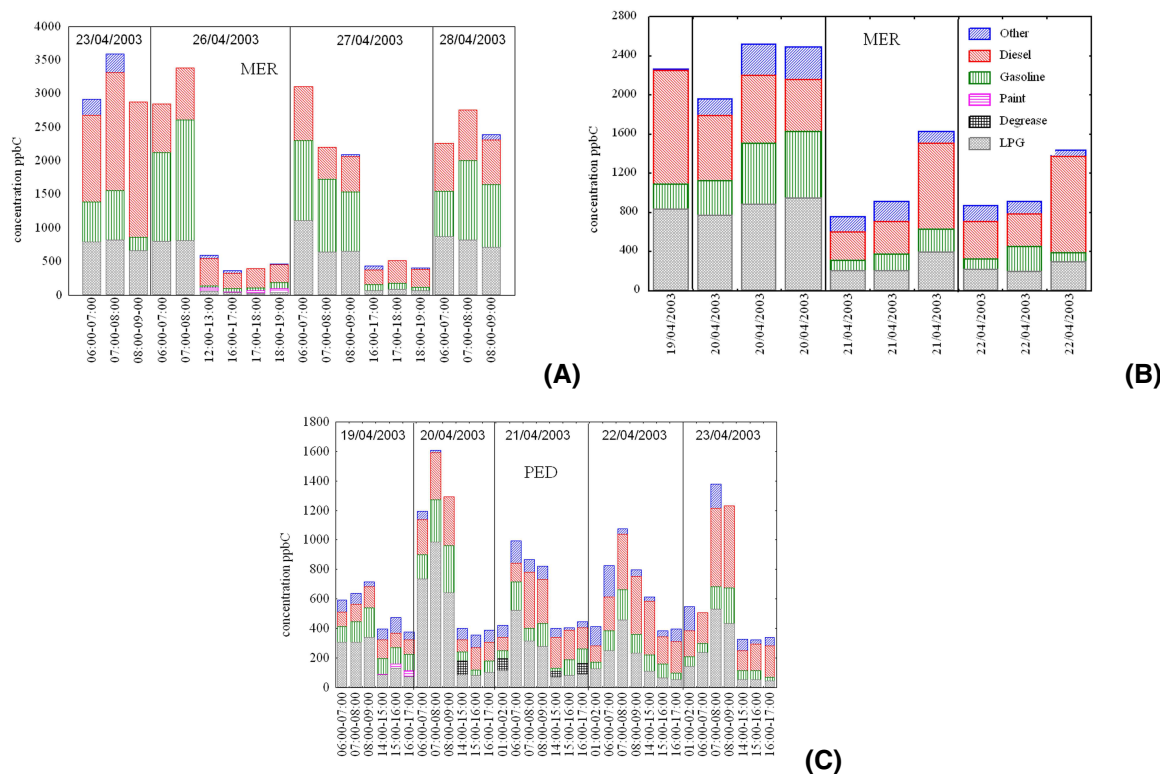
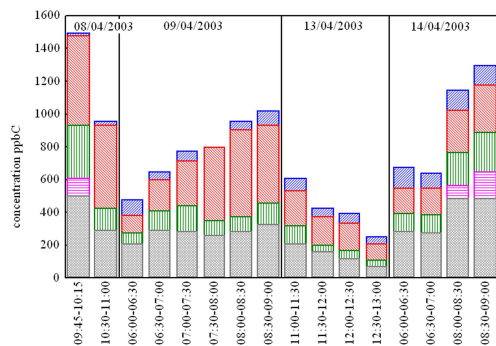


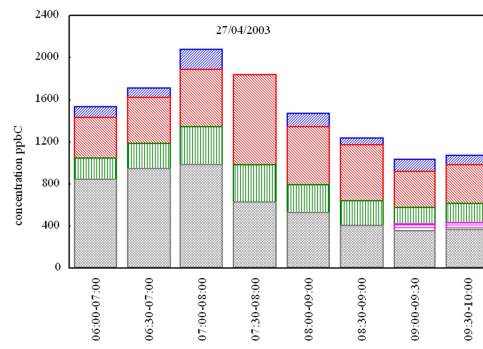
Fig. 9. (A) 60 min samples at La Merced site, (B) instantaneous samples at La Merced site; (C) 60 min samples at Pedregal site.

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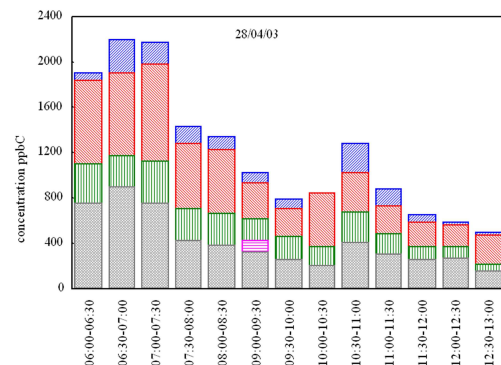
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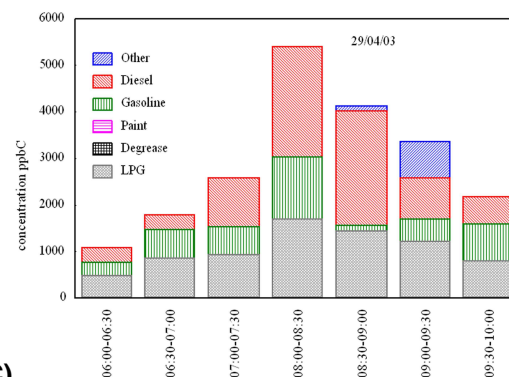
(A)



(B)



(C)



(D)

Fig. 10. Source apportionment distribution for NMHC during the 2003 campaign.

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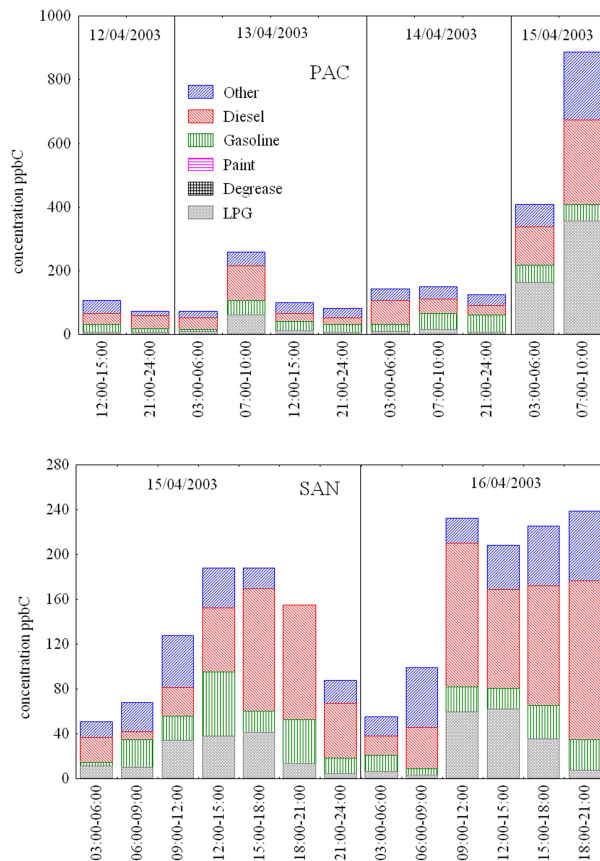


Fig. 11. Source apportionment of NMHC at two rural sites and different sampling.

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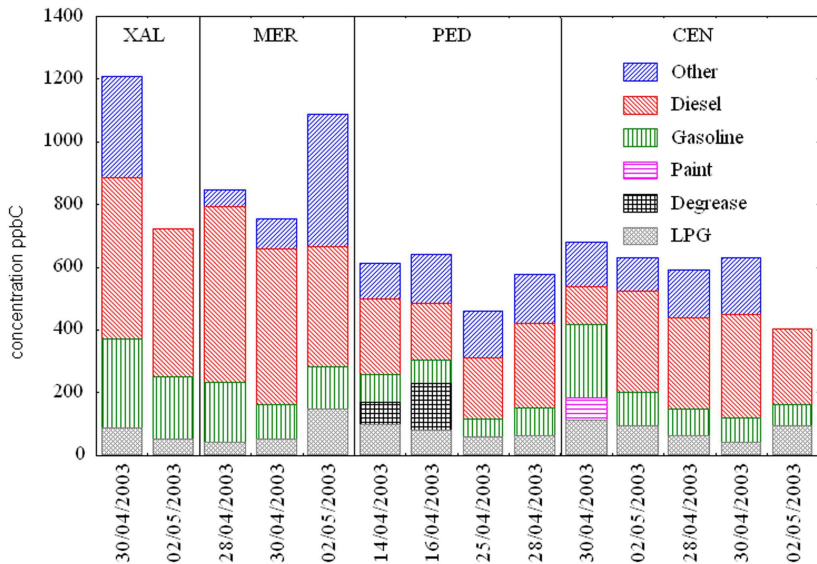


Fig. 12. Source apportionment distribution for NMHC at Xalostoc, La Merced, Pedregal and CENICA sites during the 2003 campaign 12:00–15:00 hr.



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